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- (54) BATTERY CONTAINING BIS(PERFLUOROALKYLSULFONYL)IMIDE OR CYCLIC PERFLUOROALKYLENE DISULFONYLIMIDE SALTS

BIS(PERFLUOROALKYLSULFONYL)IMID-ODER ZYKLISCHE
PERFLUOROALKYLENEDISULFONYLIMID-SALZE ENTHALTENDE BATTERIE

BATTERIE CONTENANT DES SELS BIS(PERFLUOROALKYLSULFONYL)-IMIDE ET (PERFLUOROALKYLENEDISULFONYL)-IMIDE CYCLIQUE

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Description

[0001] This invention relates to fluorinated anions of lithium salts useful in battery electrolyte compositions.

Electrolyte salts for use in electrochemical cells, e.g., lithium or lithium ion batteries, must exhibit good ionic conductivity and electrochemical, thermal, and chemical stability. In addition, the components of the electrochemical cell must be stable towards the electrolyte. Stability concerns are particularly acute in the case of electrochemical cells having aluminum cathode current collectors because aluminum is susceptible to corrosion.

[0003] Among the known electrolyte salts, lithium bis(trifluoromethanesulfonyl)imide ((CF₃SO₂)₂N·Li+) has good conductivity and stability, but is highly corrosive toward aluminum at potentials above 3V(vs Li/Li+). LiPF₆ has excellent conductivity and is noncorrosive, but is thermally and hydrolytically unstable. LiO3SCF3 (also called lithium triflate) has good thermal and chemical stability, but has low conductivity and is also highly corrosive toward aluminum at positive electrode potentials above 3V(vs Li/Li+).

[0004] Indeed, the corrosion of aluminum at potentials above 3V in electrolytes containing lithium triflate or lithium bis(trifluoromethanesulfonyl)imide is so severe as to make these salts of little use for applications in the more advanced, high voltage cells, especially rechargeable cells. Thus, the use of presently-available electrolyte salts in high voltage lithium or lithium-ion cells has resulted in batteries having suboptimal performance characteristics such as restricted operating temperature ranges, limited discharge/charge rates and inadequate cycling performance, particularly when aluminum components are used.

[0005] The present invention relates to the discovery of certain salts of fluorochemical anions that are highly conductive in nonaqueous, polar organic media and inhibit corrosion of aluminum at high oxidation potentials. These salts are therefore useful as electrolytes in high voltage electrochemical cells, such as lithium batteries, which contain aluminum components. The fluorochemical salts are comprised of a cationic portion, which is typically a metal cation and an anionic portion. The invention provides electrochemical cells or batteries containing such electrolyte salts and aluminum components.

[0006] In one aspect, the invention features a battery that includes at least one positive electrode; at least one negative electrode; at least one aluminum current collector; and an electrolyte composition comprising a salt disposed in a matrix, said salt having the formula

$$[(\mathsf{R}_{\mathsf{n}}\mathsf{SO}_2)(\mathsf{R}_{\mathsf{n}}\mathsf{SO}_2)\mathsf{N}]_{\mathsf{n}}\mathsf{M} \tag{I}$$

or

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35 (II)

in which

R₁₁ and R₁₂ are each independently a straight or branched perfluoroalkyl group of 1 to 12 carbon atoms, a perfluorocycloalkyl group or a perfluorocycloalkyl-perfluoroalkyl group of 4-7 ring carbon atoms and 1-4 carbon atoms in the alkyl chain wherein the ring carbon atoms may be optionally substituted by a straight or branched perfluoroalkyl group of 1 to 12 carbon atoms, with Rn and Rg, taken together, having a total of at least 3 carbon atoms, preferably at least 4 carbon atoms; R₁₃ is a perfluoroalkylene moiety of2 to 6 (preferably 3 to 5) carbon atoms, optionally substituted by a straight or branched perfluoroalkyl group of 1 to 12 carbon atoms or a perfluorocycloalkyl group of 4-7 carbon atoms:

M is a cation with a valence equal to n; and n is an integer of 1 to 4;

wherein the voltage, measured at the positive electrode is greater than 3.0 volts vs. Li/Li+.

[0007] In a preferred embodiment the battery has a cycle life greater than 50 cycles.

[0008] The inventive battery exhibits unexpected benefits in electrochemical performance. Specifically, the invention provides the use of electrolyte compositions that exhibit high ionic conductivity and excellent electrochemical, thermal,

and hydrolytic stability, yet at the same time inhibit degradation of aluminum battery components (e.g., corrosion of aluminum or aluminum coated components such as current collectors) at voltages typically encountered during battery operation (e.g., positive electrode (cathode) potentials greater than 3V, preferably greater than 3.5V and up to but not limited to 4 5V vs. Li/Li+). High voltage secondary batteries of this invention having aluminum components, such as the current collector, have a cycle life of greater than 50, preferably greater than 100 cycles.

[0009] The salts employed in the invention provide all of the desirable features previously associated with Li⁺N (SO₂CF₃)₂, such as high ionic conductivity and high electrochemical, thermal, and chemical stability. Furthermore, they avoid the use of toxic elements (such as As and Sb) which could be harmful to the environment, and they pose no known explosion hazard (as with perchlorate). Therefore, the salts used in this invention provide much improved properties in honaqueous electrolytes for high voltage, primary or secondary, lithium or lithium-ion batteries that contain aluminum or aluminum coated components.

[0010] Figure 1 is a cut-away view of a Li-ion battery of this invention.

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[0011] Figure 2 is a graph illustrating cycling performance of various electrolytes in the Li-ion battery of Fig. 1 by measuring battery capacity vs number of cycles.

5 [0012] Figures 3 through 5 are graphs of data plotting the potential (in volts vs. Li/Li+) of a battery test cell versus current density

[0013] Enroughout this application the following definitions apply:

[0014] "Matrix" refers to a medium (e.g., a solid, liquid, gel or plasticized polymer) in which salts according to formulae (f) and (f) may be dissolved or dispersed to form an ionically conductive electrolyte composition.

[0015] "Macromolecular material" refers to a homopolymer, copolymer, or combination thereof, which may or may not be cross-linked and/or plasticized.

[0016] "Get refers to a physically or chemically cross-linked polymer swollen with solvent.

[0017] "Battery" includes all electrical energy storage devices, including capacitors, electrochromic devices, and electrochemical cells

25 [0018] Voltages specified refer to potentials of a positive electrode measured relative to a Li/Li¹ reference electrode, except where otherwise noted.

[0019] *Oycollife* refers to the number of charge/discharge cycles measured at a minimum of 80% depth of discharge which the cattery provides without losing more than 50% of original capacity (in mAh (milliampere-hour)).

[0020] Electrosyte compositions contained in the battery of the invention include a matrix in which is disposed one contained in the battery of the invention include a matrix in which is disposed one contained in the battery of the invention include a matrix in which is disposed one contained as the structure set forth the contained and of invention, above. The electrolyte compositions based upon these salts are particularly useful in the contained as the containing aluminum (positive electrodary (rechargeable) batteries (especially secondary lithium batteries) containing aluminum (positive electrodary as the containing aluminum corrosion at voltages typically encounted as the containing aluminum (e.g., in the range from about 3.0 to 4.5V measured at the positive electrode vs. Li/Li+). [CC21] Head of the containing aluminum, the electrolyte compositions based upon these salts further provide the containing aluminum (positive electrode vs. Li/Li+). (CC21) Head of the containing aluminum (positive electrode vs. Li/Li+). (CC21) Head of the containing aluminum (positive electrode vs. Li/Li+). (CC21) Head of the containing aluminum (positive electrode vs. Li/Li+). (CC21) Head of the containing aluminum (positive electrode vs. Li/Li+). (CC21) Head of the containing aluminum (positive electrode vs. Li/Li+). (CC21) Head of the containing aluminum (positive electrode vs. Li/Li+). (CC21) Head of the containing aluminum (positive electrode vs. Li/Li+).

10022] The area saits are those of the formula

$$[(R_{11}SO_2)(R_{12}SO_2)N]_n M$$
 (I)

Answer R_{12} are each independently a straight or branched perfluoroalkyl group of 1 to 8 carbon atoms, more preferably at 15 4 carbon atoms, with R_{f1} and R_{f2} , taken together, having a total of at least 3 carbon atoms; M is a carbon R_{f1} and R_{f2} taken together, having a total of at least 3 carbon atoms; M is a carbon R_{f1} and R_{f2} taken together have at least 4.0 atoms and M is an alkali or alkaline earth metal, a transition metal, a rare earth metal, or a Group IIB or IIIB metal. [0023]

$$\begin{bmatrix} SO_2 \\ R_{f3} \\ SO_2 \end{bmatrix}_n M$$

in which B_{15} is a perfluoroalkylene group of 2 to 5 carbon atoms optionally substituted by a straight or branched per-

fluoroalkyl group of 1 to 4 carbon atoms and M and n are as defined above. More preferably, R_{13} is a perfluoroalkylene group of 3 or 4 carbon atoms. Most preferably, R_{13} has 3 carbon atoms.

[0024] Representative imide anions of Formula I are $N(SO_2C_2F_5)_2$, $N(SO_2C_2F_5)(SO_2CF_3)$, $N(SO_2C_3F_7)_2$, $N(SO_2C_3F_7)_3$, $N(SO_2C_3F_7)_4$, $N(SO_2C_3F_7)_5$, $N(SO_2C_3F_7)_6$, $N(SO_2C_3F_7)_6$, $N(SO_2C_3F_7)_6$, $N(SO_2C_3F_7)_6$, $N(SO_2C_3F_7)_7$, $N(SO_2C_$

[0025] In general, the above described bis(perfluoroalkylsulfonyl)imide and cyclic perfluoroalkylenedisulfonylimide salts can be prepared from the reaction of fluoroalkylsulfonyl fluorides, R_1SO_2F , or perfluoroalkylenedisulfonyl fluoride, $FSO_2R_1SO_2F$, with anhydrous ammonia. Symmetrical imides in which R_{11} and R_{12} are the same can be prepared in a single step using a weakly basic organic solvent such as triethylamine as shown in Scheme I, whereas unsymmetrical imides in which R_{11} and R_{12} are different must be prepared in two steps as shown in Scheme II.

SCHEME I

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SCHEME II

 $Ether $$ R_0SO_2F + NH_3 ------> NH_4^+ NH(SO_2R_0) + NH_4^+F $$ Et_3N$$ $$ R_0SO_2NH_2 + R_2SO_2F ------> Et_3NH^+ N(SO_2R_0)(SO_2R_2) + Et_3NH^+F $$$

[0026] Cyclic perfluoroalkylenedisulfonylimide salts can be prepared as described in U.S. Patent No. 4,387,222.

[0027] Perfluoroalkylsulfonylfluorides and perfluoroalkylenedisulfonylfluorides used as precursors to the imide salts of this invention can be prepared by a variety of methods known in the art as described, for example, in U.S. Patents 3,542,864; 5,318,674; 3,423,299; 3,951,762; 3,623,963; 2,732,398 and S. Temple, <u>J.Org. Chem.</u>, 33(1), 344 (1968), D.D. DesMarteau, <u>Inorg. Chem.</u>, 32, 5007 (1993).

[0028] To form the electrolyte composition, the salt is mixed with the matrix material such that the salt is at least partially dissolved or dispersed in the matrix material. The salt is preferably employed at a concentration such that the conductivity of the electrolyte solution is at or near its maximum value, although a wide range of other concentrations will also serve.

[0029] The matrix material may be in the form of a solid, liquid, gel or a liquid impregnated porous membrane. For battery applications, the matrix is chosen to provide the particular conductance, viscosity, mechanical strength, and reactivity properties desired for the electrolyte.

[0030] Although lithium (Li*) is a preferred cation M, other useful metal cations are: Na*, K*, Ca²*, Mg²*, Zn²*, and Al³*. Preferred metal cations and preferred solvents or matrix materials are dependent on cathode and anode construction in the battery. For lithium or lithium-ion batteries (having a lithium metal or lithium-ion intercalated anode) the preferred cation is Li*, and the preferred solvents are aprotic (e.g., excluding water and alcohols).

[0031] Mixtures of matrix materials can be employed and are sometimes preferred in tailoring the matrix material's properties to provide optimum performance. In general, the amount of matrix material is selected such that the salt concentration ranges from about 0.1M to about 2.0M, preferably about 1M.

[0032] Suitable matrix materials for preparing electrolyte solutions can be liquid, polymeric or mixtures of polymer and liquid. Examples of suitable solid matrix materials include polymers and copolymers such as polyethers like poly

(ethylene oxide), polyesters, polyacrylates, polyphosphazenes, polysiloxanes, poly(propylene oxide), fluoropolymers (e.g., poly(vinylidene fluoride)), and poly(acrylonitrile), as well as the polymers and copolymers described in Armand et al., U.S. Patent No. 4,505,997, and mixtures thereof. The polymers may be used in cross-linked or uncross-linked form and or plasticized. Such materials are generally dry, i.e., have a water content less than about 100 ppm, preferably less than about 50 ppm.

[0033] In batteries comprising a highly reducing electrode (such as lithium metal) and a liquid matrix material, the liquid is preferably a nonaqueous, polar, aprotic, organic solvent. Such liquids are generally dry, i.e., have a water content less than about 100 ppm, preferably less than about 50 ppm. Examples of suitable aprotic liquids include linear ethers such as diethyl ether, diethylene glycol dimethyl ether, and 1,2-dimethoxyethane; cyclic ethers such as tetrahydrofuran. 2-methyltetrahydrofuran, dioxane, dioxolane, and 4-methyldioxolane; esters such as methyl formate, ethyl formate, methyl acctate, dimethyl carbonate, diethyl carbonate, propylene carbonate, ethylene carbonate, and butyrolactones (e.g., gamma butyrolactone); nitriles such as acetonitrile and benzonitrile; nitro compounds such as nitromethane or nitrobenzene; amides such as N,N-dimethylformamide, N,N-diethylformamide, and N-methylpyrrolidinone sulfoxides such as dimethyl sulfoxide; sulfones such as dimethylsulfone, tetramethylene sulfone, and other sulfolianes exazolidinones such as N-methyl-2-oxazolidinone and mixtures thereof. Maximum conductivities of the electrolyte salts of this invention in typical nonaqueous, polar, aprotic liquid media (e.g., propylene carbonate) are generally in the range of 0.1 - 20 mS, (milliSiemens) at room temperature, preferably greater than 1 mS.

[0034] In some cases, it may be desired to add other salts to the electrolyte composition in order to maximize performance such as corrosion properties, conductivity of battery components by the electrolyte). Such salts include, but are not limited to, alkali metal, alkaline earth metal, and Group IIIB metal (e.g., aluminum) salts of anions such as NO_3^+ , $BF_4^- BF_5^-$, CIO_4^+ , SbF_6^+ , $R_1SO_3^-$ (in which R_1 is a perfluoroalkyl group having between 1 (preferably 2) and 12 carbon atoms inclusive); a bis-(perfluoromethylsulfonyl)imide anion; an anion having a formula selected from the group consisting of $F_{11}R_{12}N$ -(CF_2)_n- SO_2 - X^-

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$$Z = N - (CF_2)_{\overline{n}} - SO_2 - X$$

nian this Color N'SOARm, or

$$(R_{f5} SO_2) - C - (SO_2 R_{f4});$$

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or SE_1 and B_2 independently, are $-CF_3$, $-C_mF_{2m+1}$, or $-(CF_2)_q$ - SO_2 - X^*M^+ ; B_{13} , B_{14} , and B_{15} , independently, are $-SE_1$ and $-SE_2$ and $-SE_3$ are $-SE_3$ and $-SE_3$ $-SE_3$ are $-SE_3$ are $-SE_3$ and $-SE_3$ are $-SE_3$ are $-SE_3$ are $-SE_3$ are $-SE_3$ and $-SE_3$ are $-SE_3$ are $-SE_3$ are $-SE_3$ and $-SE_3$ are $-SE_3$ are $-SE_3$ are $-SE_3$ are $-SE_3$ are $-SE_3$ and $-SE_3$ are $-SE_3$ and $-SE_3$ are $-SE_3$ are $-SE_3$ are $-SE_3$ and $-SE_3$ are $-SE_3$ are $-SE_3$ and $-SE_3$ are $-SE_3$ and $-SE_3$ are $-SE_3$ and $-SE_3$ are $-SE_3$ are $-SE_3$ are $-SE_3$ and $-SE_3$ are $-SE_3$ are

$$N - (CF_2)_{\overline{n}}$$

$$R_{f2}$$

or

$$Z$$
 R_{f7}
 $N-(CF_2)_{\overline{n}}$

 R_{16} is $-CF_3$, $-C_mF_{2m+1}$, or $-(CF_2)_q$ - SO_2 -X-M+; R_{16} and R_{17} , independently, are perfluoroalkylene moieties having the formula $-C_rF_{2r}$; n is 1-4; r is 1-4; r is 1-12 preferably 1-8; and q is 1-4; (such salts are described by Waddell, et al. in U.S 5514493); a bis-perfluoroalkylsulfonyl methide anion R_rSO_2 - $C^*(R)$ - SO_2 - R_1^* in which R_1 and R_1^* , independently, are perfluoroalkyl groups having between 1 and 12 carbon atoms, inclusive, and R_1 is R_1 , R_2 , and R_3 , independently, are perfluoroalkylsulfonyl) methide anion of the formula $-C(SO_2R_1)(SO_2R_1^*)(SO_2R_1^*)$ in which R_1 , and R_2 , independently, are perfluoroalkyl groups having between 1 and 12 carbon atoms, inclusive.

[0035] Representative examples of suitable salts include, LiNO3, LiBF4, LiAsF6, LiClO4, LiPF6, CF3SO3Li, C2F5SO3Li, C10F21SO3Li, (CF3SO2)2NLi, (CF3SO2)2NNa, [(CF3SO2)2N]3Al, (CF3)2NC2F4SO3Li, (CF3SO2)2C(CH3)Li, cyclo(CF2SO2)2C(C6H5)Li, (CF3SO2)3CLi, ((CF3)2NC2F4SO2)2NLi and mixtures thereof.

[0036] Preferred salts are nitrate salts (NO₃*). Addition of nitrate salts to the electrolyte composition has been found to provide improved uniformity of current density at relatively higher temperatures. The counterion of the nitrate salt can be any counterion that will provide sufficient ionization within the electrolyte solution to provide useful conductivity properties. Nitrate salts that are preferred include nitrate salts having counterions such as: a tetraalkyl ammonium ion (NR₄, wherein each R can independently be a lower alkyl such as methyl, ethyl, butyl, etc.); an alkali metal; an alkaline earth metal; a Group IIB metal; a Group IIB metal; or mixtures thereof.

[0037] Addition of small amounts of these salts to the electrolyte composition is preferred because these salts have been found to reduce corrosion and prevent increases in cell resistance in batteries operated at, or subjected to, elevated temperatures. The salts may be added in any amount that provides such results, with concentration from about 0.1 mM (millimolar) to about 500 mM, preferably from about 0.1 mM to about 100 mM, most preferably between 1.0 mM and 50 mM. When used in relatively low concentrations the nature of the cation is not critical for many applications, although lithium (i.e. LiClO₄) is especially preferred.

[0038] The anodes and cathodes of the inventive electrochemical cell are generally particles of active material blended with a conductive diluent such as carbon black and graphite and bound into a plastic material. Typical binders are polyvinylidene fluoride, ethylene-propylene-diene (EPDM) terpolymer, and emulsified styrenebutadiene rubber (SBR) among others, and the binder may be cross-linked. The binder may be for example a solid carbon matrix formed from the thermal decomposition of an organic compound. The composite electrode material is in turn applied to an expanded metal screen or metal foil (preferably aluminum) current collector using a variety of processes such as coating, casting, pressing, or extrusion. In polymer electrolyte batteries the polymer electrolyte can act as the active material binder. [0039] Some examples of negative electrodes are carbon based materials such as graphite, coke, carbon fiber, and pitch, transition metal oxides such as LiTi₅O₁₂ and LiWO₂, lithium metal and lithium alloys. In the case of lithium-ion batteries, the lithium may be intercalated into a host material such as carbon, (i.e., lithiated carbon) or carbon alloyed with other elements such as Si, B and N, a conductive polymer or an inorganic host material that is intercalatable, such as Li_xTi₅O₁₂. Li_xV₂O₅, Li_xV₆O₁₃, and Li_xMn₂O₄ and Li_xMnO₂ (the spinel and alpha forms respectively). The material comprising the anode (negative electrode) of an electrochemical cell may be carried on foil (e.g., nickel or copper) backing or pressed into expanded metal screen and alloyed with various other metals.

[0040] Active cathode (positive electrode) materials provide cell voltages of at least 3.0 volts at a full state of charge relative to Li/Li⁺ and include, but are not limited to Li_xCoO₂, Li_xNiO₂, Li_xMn₂O₄ and Li_xMnO₂, V₂O₅, V₆O₁₃, and fluorinated carbon, including the charged and discharged forms of these materials, and conducting polymers such as polypyrrole and polyvinylferrocene.

[0041] The invention includes primary and secondary batteries. In primary batteries, the cathode (positive electrode) could be fluorinated carbon (CF_x)_n, SO₂, SO₂CI₂, or Ag₂ CrO₄.

[0042] The invention is illustrated further by the following examples.

EXAMPLES

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[0043] The salts Li+*N(SO₂CF₃)₂ and Li+*OSO₂CF₃ used in the examples are commercially available in high purity from Minnesota Mining and Manufacturing Company as Fluorad™ Lithium Trifluoromethanesulfonimide Battery Electrolyte, HQ-115 and as Fluorad™ Lithium Trifluoromethanesulfonate, FC-122, respectively. High purity, battery grade Li+PF₆* was purchased from Hashimoto Chemical Co., Ltd. through a US distributor (Biesterfeld U.S., Inc.). All salt samples were stored and handled in a nitrogen or argon filled dry box (Vacuum Atmospheres Inc.) to prevent contam-

ination by water.

Example 1

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Lithium (trifluoromethanesulfonyl) (perfluorobutanesulfonyl)imide

[0044] Under a nitrogen atmosphere, a dry, 500 mL Fisher-Porter pressure bottle equipped with a stainless steel pressure head and magnetic stirrer was charged with 35.00 g CF₃SO₂NH₂ (prepared, e.g., as described in Foropoulos and DesMarteau. Inorg. Chem., 23:3720-23 (1984)), 98 mL anhydrous triethylamine and 74.55 g C₄F₉SO₂F. The pressure vessel was sealed and the reaction mixture heated to 90°C with stirring for 17 hours. The volatile components of the reaction mixture were evaporated under reduced pressure at 70°C then the reaction mixture was treated with a mixture of 700 mL water and 700 mL methylene chloride with stirring to form a two phase mixture. The methylene chloride phase was then separated, washed with two 700 mL portions of water, dried over anhydrous MgSO4, filtered and then evaporated under reduced pressure to yield a dark red liquid. This liquid was combined with 500 g of polyphosphoric acid (Aldrich Chemical Co.) in a short path vacuum distillation apparatus equipped with an ambient air-cooled condenser, and then vacuum distilled at ca. 0.1 Torr. The distillate (96.5 g, corresponding to HN(SO₂CF₃)(SO₂C₄F₉), which solidified in receiver at 0°C, was further purified by sublimation at 70°C, 1.0 Torr and then added in portions to a sturry of 52 g of Li₂CO₃ (99.997% purity, from Aldrich Chemical Co.) in 800 mL of methyl t-butyl ether with stirring at room temperature. After ca. 1 hour CO₂ evolution subsided and the reaction mixture was heated to ca. 40°C for 3.5 hours to complete the neutralization. The mixture was filtered by gravity through filter paper and then filtered again by suction through a 0-22 micron Tefsep™ Teflon membrane (Micron Separations Inc.) to remove particulates. The filtrate was evaporated under reduced pressure at 25-80°C to yield a clear, colorless oil. Two 250 mL portions of toluene were combined with the oil and the mixture evaporated at 40-70°C, 20 Torr after each toluene addition, causing the oil to eventually solidify giving a fine white powder. The solid was transferred to a glass jar and dried in vacuo at 100°C, 10°2 Torritor 24 hours to yield 77.0 g of the anhydrous title salt. The structure of the product was confirmed by ¹H and ¹⁹F NMR coachescopy which indicated that the purity of the salt was greater than 99% by weight.

Example 3

Little in the Aramethanesulfonyl) (perfluoroethanesulfonyl)imide

[0045] The restriction atmosphere, a dry 3L flask equipped with a dry ice condenser, mechanical stirrer, and a spanding time was charged with 500 mL anhydrous methyl t-butyl ether (MTBE) and cooled below 0°C in a dry ice bath To the region a solvent was added 500 g of crude CF₃CF₂SO₂F mixture (containing ca. 23% CF₃CF₂SO₂F by weight in 30 control of the sevent). Gaseous anhydrous ammonia (194 g) was gradually introduced through the sparging the resulting mixture at ca. 0°C with vigorous stirring. Once ammonia addition was complete, the reaction was a was removed and the reaction solution GREEN LAND TO The temperature with stirring. Excess ammonia was allowed to evaporate while stirring overnight at a transfer of the reaction solution was cooled again to 0°C and treated with 83.9 g LiOH-H₂O dissolved in 750 m. If we for with stirring. The mixture was filtered through filter paper by gravity to remove LiF precipitate. To the filtrate was accommentated aqueous HCI in portions with stirring at 0°C until the pH of the mixture was 0. Ling two-phase mixture was separated into separate aqueous and ether phases. The aqueous phase was extracted with two additional 400 mL portions of MTBE. The ether fractions were combined, extracted with two 500 multiprishes of water, and then dried over anhydrous MgSO4 overnight. The ether solution was filtered by gravity through the paper and then by suction through a 0.22 micron Tefsep™ membrane (Micron Separations Inc.). The filtrate was concentrated by rotary evaporation, hexane was added, and the solution concentrated again. This was repeated into the product crystallized from solution as a white solid. The product (Crop 1) was isolated by suction filtration in rough a glass frit giving 33.0 g. A second crop of crystals (20.2 g) was recovered in the same manner from the titrate by concentrating further followed by filtration. The remaining filtrate was evaporated to dryness at 40°C, 20 mm to visit a third crop of relatively crude product (26.8 g). All product fractions (Crops 1-3) were dried at 10°2 Torr at room temperature for call 15 min to give a combined yield of 80.0 g. Quantitative ¹H and ¹³C NMR analysis of the product from Disp 2 indicated that it was the desired product with a purity of 98% by weight.

[0047] Under a hitrogen atmosphere, a dry Parr 4560-Series Benchtop Mini Reactor equipped with a 600 mL capacity Money reactor cylinder, mechanical stirrer, thermocouple and heating mantle was charged with 53 g CF₃CF₂SO₂NH₂ (Cross 1 - 2) and 152 mL anhydrous triethylamine. While cooling the reactor at -78°C in dry ice, 55.2 g of gaseous CF₃SO₂F an odes purity) was condensed into the reactor with stirring. The reactor was sealed and the temperature of the react on mixture gradually raised to 90°C with vigorous stirring, then held at 90°C with stirring for a total of 24 hours. The volatile components of the reaction mixture were evaporated under reduced pressure at 70°C, then the

residue was treated with a mixture of 700 mL water and 700 mL methylene chloride with stirring to form a two phase mixture. The methylene chloride phase was then separated, washed with two 700 mL portions of water, dried over anhydrous MgSO₄, filtered and then evaporated under reduced pressure to yield a dark red liquid. This liquid was combined with 600 g of polyphosphoric acid (Aldrich Chemical Co.) in a short path vacuum distillation apparatus equipped with an ambient air-cooled condenser, and then vacuum distilled at ca. 15 Torr, 85-88°C. The distillate (78.1 g, corresponding to $HN(SO_2C_5F_3)(SO_2C_5F_5)$), which solidified in receiver at 0°C, was converted to the corresponding lithium salt, and further purified, using essentially the procedure described in Example 1 to yield 75.0 g of the anhydrous title salt. The structure of the product was confirmed by 1H and ^{19}F NMR spectroscopy which indicated that the purity of the salt was 98% by weight.

Example 3

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Lithium Bis(perfluoroethanesulfonyl)imide

[0048] As in Example 2, a dry Parr 4560-Series Benchtop Mini Reactor was charged with 155 mL anhydrous triethylamine. While cooling the reactor at -78°C in dry ice, gaseous C₂F₅SO₂F (100.0 g, > 99% purity) followed by anhydrous ammonia (3.51 g) were condensed into the reactor with stirring. The reactor was sealed and the temperature of the reaction mixture gradually raised to 90°C with vigorous stirring. The reaction mixture was held at 90°C with stirring for a total of 20.5 hours. The intermediate product, HN(SO₂C₂F₅)₂, was isolated as previously described, then added to a slurry of Li₂CO₃ (45.7 g, 99.997% purity) in 800 mL of methyl t-butyl ether to produce the lithium salt. This crude lithium salt was further purified using essentially the procedure described in Example 1 to yield 73.34 g (92% yield based upon NH₃) of the anhydrous title salt. The structure of the product was confirmed by ¹H and ¹9F NMR spectroscopy which indicated that the purity of the salt was 99.9% by weight.

25 Example 4

Lithium Bis(perfluorobutanesulfonyl)imide

[0049] As in Example 1, a dry, 500 mL Fisher-Porter pressure bottle was charged with anhydrous triethylamine (100.5 g) and C₄F₉SO₂F (145.0 g). The pressure vessel was sealed, cooled to -78°C in dry ice and then charged with of anhydrous ammonia (3.0 g) with stirring. After warming gradually to room temperature, the reaction mixture was heated to 90°C with stirring for 24 hours. The intermediate product, HN(SO₂C₄F₉)₂, was isolated as previously described, then added to a slurry of Li₂CO₃ (40.0 g, 99.997% purity) in 800 mL of methyl t-butyl ether to produce the lithium salt. This crude lithium salt was further purified using essentially the procedure described in Example 1 to yield 90.8 g (86% yield based upon NH₃) of the anhydrous title salt. The structure of the product was confirmed by ¹H and ¹9F NMR spectroscopy which indicated that the purity of the salt was 98% by weight.

Example 5

40 Lithium Bis(perfluoropropanesulfonyl)imide

[0050] As in Example 4, the title compound was prepared using anhydrous triethylamine (100.5 g), C₃F₇SO₂F (100.0 g, 68:32 *iso:normal* isomer ratio), and anhydrous ammonia (3.0 g). A total of 40.8 g of the anhydrous title salt was recovered as a light pink solid. Quantitative analysis by ¹H and ¹⁹F NMR spectroscopy indicated that the product contained the following major components in order of decreasing weight-%: Li⁺N(SO₂C₃F₇)₂, 85.1%, 39:61 iso:normal C₃F₇ ratio; Li⁺N(SO₂i-C₃F₇)(SO₂F), 8.7%; Li⁺N(SO₂n-C₃F₇)(SO₂F), 4.2%.

Example 6

50 Lithium cyclic-1,3-perfluoropropanedisulfonylimide

[0051] The acidic form of the title compound, HN(SO₂C₃F₆SO₂), was prepared according to the method described in Koshar, US Patent 4,387,222, Example 1 and converted to the anhydrous lithium salt using the method described in Example 4 of the present description. A total of 26.4 g (56% yield, based upon 1,3-perfluoropropanedisulfonylfluoride) of the anhydrous title salt was isolated with a purity of 98.7% by ¹H and ¹⁹F NMR spectroscopy.

Example 7

Lithium cyclic-1,2-perfluoroethanedisulfonylimide,

[0052] The acidic form of the title compound, HN(SO₂C₂F₄SO₂), was prepared according to the method described in Koshar. US Patent 4,387,222, Example 3 and converted to the anhydrous lithium salt using the method described in Example 4 of the present description. A total of 16.6g (61% yield, based upon 1,2-perfluoroethanedisulfonylfluoride) of the anhydrous title salt was isolated with a purity of 99.8% by ¹H and ¹⁹F NMR spectroscopy.

[0053] The inventive electrolyte compositions are particularly useful in that they control or prevent aluminum corrosion in electrochemical cells operating at voltages greater than 3V and up to but not limited to 4.5V, referenced to lithium metal.

[0054] One way to assess the extent of aluminum corrosion in the presence of a particular electrolyte solution is to measure the anodic current density vs. time response at a fixed d.c. potential of a cell containing an aluminum electrode having a freshly exposed aluminum surface, as described in the Examples section below. The higher the current density, the faster aluminum corrosion is occurring.

[0055] Another practical measure of the compatibility of a particular electrolyte solution with aluminum is given by the cycling performance of a high voltage, secondary (or rechargeable) battery containing the electrolyte solution of interest and a cathode comprising an aluminum current collector.

20 Test Methods

[0056] Corresion current measurements were made according to the technique generally described in Bard and Faulkner Electrochemical Methods: Fundamentals and Applications, John Wiley and Sons, New York, 1980, pp. 350-353. An electrochemical cell was constructed having an aluminum working electrode. A lithium wire inserted in a luggin capitary (i.e., a glass capillary into which the electrode was inserted) served as a reference electrode and a 10 cm² diatinum flag was used as an auxiliary electrode. The working electrode was fabricated from a 99.999% aluminum rod inserted into a Teffon sleeve to provide a planar electrode area of 0.07 cm². The native oxide layer was removed by 1 rst polishing the planar working surface with 3 \(\mu\) m aluminum oxide paper using hexane as a lubricant under ambient conditions. To lowed by a second polishing under an argon atmosphere.

[0057] After coisning, the cell was assembled under argon, connecting the three electrodes to a potentiostat. The solvent risd been previously dried by stirring over CaH2 for 24 hrs., followed by vacuum distillation. The electrolyte sait was dried under vacuum at 120°C for at least 18 hrs. The final water content of the electrolyte solution was less than 10 ccm. is measured by Karl Fischer titration. The aluminum electrode was polarized at +4.2V vs. Li/Li+ (iR 10 ccm. 1.1.2 ccm.

[0058] Consider the asurements (in microamps/cm²), taken after one hour, are summarized in Table I, below. Additionally and a surger the current vs. time curve (millicoulombs, mC) is used as a measure of the maximum possible and a maximum corresion occurring. This data is also summarized in Table I. Data for the following salts is included for run as a time curves:

Lit N 10,000, 2 Lit O₃SCF₃ and Lit PF₆ (designated COMP1, COMP2 and COMP3, respectively in Table I).

[0059] Another feature which is important to battery performance is ionic conductivity. Conductivity measurements of the machine stronger at a concentration of 1M in PC/DME, propylene carbonate/dimethoxy ethane (1:1 by volume), are also recorded in Table I.

TABLE I

	Example	Current Density μA/cm²	Integrated Chronoamperiometric Area (mC)	Conductivity 1M (mS*)		
	1	5 19	2.31	9.4		
	2	2 3	2.48	10.6		
	3	2.00	0.94	9.8		
	١ ـ ا	2.51	0.7	7.1		
	5	1 26	0.83			
	ć	2.31	. 1.37	11,1		
	Comp :	7,570	2360	9.9		
Í	Comp 2	17.600	5860	6.4		
	Camp 3	0.9	0.7	17		

mS = mill-Siemens

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[0060] The data in Table I shows that while LiN(SO₂CF₃)₂ and LiO₃SCF₃ are highly corrosive toward aluminum and produce large corrosion currents, the salts of the present invention are at least three orders of magnitude less corrosive and produce corrosion currents comparable to LiPF₆, a widely used lithium salt known to be noncorrosive when in contact with aluminum at high potentials. Additionally, the data show that salts used in the present invention provide ionic conductivities useful for high performance battery applications.

[0061] High performance can be demonstrated in cells under typical cycling conditions using high voltage, rechargeable lithium ion batteries constructed with aluminum positive current collectors and suitable electrode materials. Electrolyte salts compatible with aluminum will provide long cycle life with minimal loss in capacity; whereas, corrosive salts cause a sharp drop in battery capacity (mAh) after a relatively small number of cycles.

[0062] In Examples 8 and 9 and in Comparative Examples 4-6, batteries (as shown in Figure 1) were prepared and subjected to a minimum of fifty charge-discharge cycles using a commercial battery tester from Maccor Inc., Tulsa Oklahoma to measure the capacity of the cells. The anode 1 was prepared using a mixture of 85.5 wt% XP grade petroleum coke (Conoco Co), 4.5 wt% Ensagri Super "S" carbon black and 10 wt% polytetrafluoroethylene (PTFE) binder, mixed together in a grinding device such as a food processor or coffee grinder and pressed into pellets approximately 12 to 14 mils (.31 to .36 mm) in thickness and 7 mm in diameter. The cathode was formulated using a mixture of 83.7 wt.% LiCoO₂, 6.3 wt.% Shawinigan carbon black and 10 wt.% PTFE mixed as described above and pressed into pellets about 0.33 mm thick and 7 mm in diameter. The cell stack assembly comprised a 31 mil (.80 mm) copper disk 3 as the anode current collector, 14 mil (.36 mm) thick anode, a 2 mil (0.05 mm) porous polyethylene cell separator 5, 12.5 mil (.32 mm) thick cathode 7 of LiCoO₂ and a 20 mil (.51 mm) thick aluminum cathode current collector 9. The stack was placed into the chromium steel "1225" coin cell can 10 (12 mm in diameter and 2.5 mm stack depth) and 40 μL of 1.0 M electrolyte were added. The can was then compressively sealed with the polypropylene gasket 12 and stainless steel top 14. All cell assembly operations were done in a dry atmosphere. The batteries tested were as follows:

Example 8 1M LiN(SO₂C₂F₅)₂, Example 3, in 1:1 (volume) mixture of ethylene carbonate/dimethyl carbonate

Example 9 1M LiN(SO₂)₂(CF₂)₃, Example 6, in 1:1 (volume) mixture of ethylene carbonate/dimethyl carbonate

Comparative Example 4 1M LiN(SO₂CF₃)₂ in 1:1 (volume) propylene carbonate/dimethoxy ethane

Comparative Example 5 1M LiSO₃CF₃ in 1:1 (volume) propylene carbonate/dimethoxy ethane

Comparative Example 6 1M LiPF₆ in 1:1 (volume) propylene carbonate/dimethoxy ethane

[0063] Each battery was individually loaded into the battery tester and charged at a constant voltage of 4.2V, with a maximum current limitation of 5mA, until the current dropped to 0.1mA (end of charge). Each cell was then discharged under a constant current of 0.8mA until the cell voltage reached 2.75V (end of discharge). After approximately ten charge/discharge cycles each cell was charged under constant voltage of 4.2V with a maximum current limit of 5mA until the current reached 0.1mA. The cells were then discharged under constant current of 0.8mA until the cell voltage reached 3V. The current for each cell was then interrupted (0 current) and the voltage recorded for 30 minutes under zero current conditions to provide a measure of internal cell resistance. Each cell was then charged under a constant voltage of 4.2V with a maximum current limit of 5mA for a total of 24 hours. The cell was then discharged at a constant current of 0.8mA to 3.0V and a 30 minute current interrupt was applied. After the interrupt the cell is discharged to 2.75V under a constant current of 0.8mA. The procedure was repeated for a minimum of fifty cycles.

[0064] The results of these tests are shown in Figure 2 as a plot of battery capacity (mAh) vs. cycle number for each of the batteries tested.

[0065] The lines plotted are:

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line A - Example 8 LiN(SO₂C₂F₅)₂

line B - Comparative Example 6 LiPF6

line C - Example 9 LiN(SO₂)₂ (CF₂)₃

line D - Comparative Example 4 LiN(SO₂CF₃)₂

line E - Comparative Example 5 LiO₃SCF₃ (lithium triflate)

[0066] In Figure 2, the smooth portion of each line provides a measure of cell capacity as a function of cycle number under conditions of relatively rapid rate of cell discharge. The occasional spikes in Figure 2 are the result of increased capacity resulting from the application of a current interrupt preceded by a 24 hour charging period. The test procedure demonstrates the effect of a 24 hour charging period (in which corrosion is most likely to occur) on cell capacity before and after the charging period. The cycling data of Figure 2 shows that LiO₃SCF₃ and LiN(SO₂CF₃)₂ cause battery

capacity to drop precipitously to zero after just one or two cycles. The electrolyte salts of the present invention provide more than fifty cycles with less than a 20% loss in cell capacity. The data further demonstrates that the salts of the present invention provide cycling performance comparable to or exceeding LiPF₆, a salt commonly employed in commercial lithium-ion batteries.

[0067] The inventive batteries with the specified electrolyte salts are unique in that aluminum corrosion is suppressed at high electrochemical potentials (potentials greater than +3,0 Volts vs. Li/Li+), while at the same time providing very good ionic conductivity and stability (e.g., thermal, electrochemical, and hydrolytic stability). The batteries of Examples 8 and 9 showed no significant increase in internal cell resistance after the 24 hour potential hold.

[0068] As shown in Table I and Figure 2, electrolyte compositions used in the invention are less corrosive than known electrolyte compositions containing other perfluoro-organosulfonyl salts. In some instances, electrolyte compositions used in the invention approach or exceed the performance of inorganic salts such as Li* PF₆ which are known not to promote corrosion.

[0069] Examples 10-14 show the effect of adding electrolyte salts to batteries operated at elevated temperatures. In these Examples the electrochemical cell (described under Test Methods) was filled with 1.0 M lithium bis(perfluoroethylsu.fonyl)imide in 1:1 (vol) ethylene carbonate/dimethyl carbonate. The aluminum electrode was polarized for one hour at 4 2V L:/Li*, and then the potential was increased in 50mV increments while recording the current density in μΑ/cm² The added electrolyte, concentration, temperature and corresponding Figures for these Examples are shown in Table II

TABLE II Example Added electrolyte (concentration (mM)) Temperature Figure 10 None 25°C 11 None 60°C 3 12 LiNO₃ (25) 60°C 3, 4, 5 13 LiNO₃ (13) 60°C 3 14 LiNO₃ (6.5) 60°C 15 LiNO₃ (3.25) 60°C 3 16 LiSO₃CF₂CF₂N(CF₃)₂ (100) 60°C 4 17 $LiSO_3CF_2CF_2N(CF_3)_2$ (21) 60°C 4

LiCIO₄ (25)

[0070] The results of these tests are shown in Figures 3 through 5, which plot current density versus potential (in increased temperature (60°C), as indicated by Example 11. With the added electrolyte saits in Eximple 12/18 this increase in current density is suppressed at voltage levels below about 4.8V, as is indicated by the relatively constant current density below this voltage.

Claims

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1. A battery comprising

at loast one positive electrode:

it is ist one negative electrode;

at least one aluminum current collector; and

an electrolyte composition comprising a salt disposed in a matrix, said salt having the formula

BEST AVAILABLE

60°C

$$[(R_{f1}SO_2)(R_{f2}SO_2)N]_n M \text{ or } \begin{bmatrix} SO_2 \\ R_{f3} \\ SO_2 \end{bmatrix}_n M$$

in which

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 R_{f1} and R_{f2} are each independently a straight or branched perfluoroalkyl group of 1 to 12 carbon atoms, a perfluorocycloalkyl group or a perfluorocycloalkyl perfluoroalkyl group of 4-7 ring carbon atoms and 1-4 carbon atoms in the alkyl chain wherein the ring carbon atoms may be optionally substituted by a straight or branched perfluoroalkyl group of 1 to 12 carbon atoms, with R_{f1} and R_{f2} taken together having a total of at least 3 carbon atoms;

 $R_{\rm B}$ is a perfluoroalkylene moiety of 2 to 6 carbon atoms optionally substituted by a straight or branched perfluoroalkyl group of 1 to 12 carbon atoms or a perfluorocycloalkyl group of 4-7 carbon atoms; M is a cation with a valence equal to n; and

n is an integer of 1 to 4, wherein the voltage measured at the positive electrode is greater than 3.0 volts vs. Li/Li+.

- 2. The battery of claim 1 wherein M is lithium.
- The battery of claim 1 wherein said salt is lithium bis(perfluoroethanesulfonyl)imide.
 - 4. The battery of claim 1 further comprising a salt selected from alkali metal, alkaline earth metal, rare earth and Group IIB and IIIB metal salts of anions selected from NO₃"; BF₄"; PF₆"; AsF₆"; ClO₄"; SbF₆"; R_fSO₃" in which R_f is a perfluoroalkyl group having between 2 and 12 carbon atoms, inclusive, bis-(perfluoromethylsulfonyl)imide; R_{ft}R_{f2}N -(CF₂)_{n1}- SO₂-X";

$$Z = \frac{R_{f6}}{N - (CF_2)_{\overline{n_1}}} SO_2 - X$$

in which X is -O', -N'SO2R13, or

45 Z is -CF₂-, -O-,

or -SF₄-; R_{f1} and R_{f2} , independently, are -CF₃, -C_mF_{2m+1}, or -(CF₂)_q-SO₂-X^{*}M⁺; R_{f3} , R_{f4} , and R_{f5} , independently, are -CF₃, -C_mF_{2m+1}, -(CF₂)_q-SO₂-X^{*}M⁺,

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$$\begin{array}{c}
R_{f1} \\
N - (CF_2)_{\overline{n_1}}
\end{array}$$

Of 10

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$$Z = N - (CF_2)_{\overline{n_1}}$$

 R_{18} is -CF₃, -C_mF_{2m+1}, or -(CF₂)_q-SO₂-X^{*}M^{*}; R_{16} and R_{17} , independently, are perfluoroalkylene moieties having the formula -C_rF_{2r}-, n₁ is 1-4; r is 1-4; m is 1-12; and q is 1-4; bis-perfluoroalkylsulfonyl methide anions having the 20 $formula\ P_f-SO_2-C^*(R)-SO_2-R_f^*\ in\ which\ R_f\ and\ R_f^*,\ independently,\ are\ perfluoroalkyl\ groups\ having\ between\ 1\ and\ R_f^*$ 12 carbon atoms, inclusive, and R is H, Br, Cl, I, an alkyl group having between 1 and 20 carbon atoms, inclusive, aryl, or alkylaryl, and a tris(perfluoroalkylsulfonyl)methide anion of the formula -C(SO_2R_1)(SO_2R_1) in which B, B, and B, and B, and ependently, are perfluoroalkyl groups having between 1 and 12 carbon atoms, inclusive

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- 5. The battery of claim 1, further comprising LiNO₃.
- The battery of claim 1, wherein the battery is rechargeable.
- 30 7. The pattery of claim 6, wherein the battery has a cycle life greater than 50 cycles.

Patentansprüche

35 1. Batterie umfassend

mindestens eine positive Elektrode,

mindestens eine negative Elektrode,

mindestens einen Stromabnehmer aus Aluminium und

eine Elektrolytzusammensetzung, umfassend ein in einer Matrix verteiltes Salz, wobei das Salz die Formel autweist

$$[(R_{f1}SO_2)(R_{f2}SO_2)N]_n M oder \begin{bmatrix} SO_2 \\ R_{f3} \end{bmatrix} M$$

in der

ក_ក und R_R jeweils unabhängig voneinander einen gerad- oder verzweigtkettigen Perfluoralkylrest mit 1 bis 12 Kohlenstoffatomen, einen Perfluorcycloalkylrest oder einen Perfluorcycloalkylperfluoralkylrest mit 4-7 Ringkohlenstoffatomen und 1-4 Kohlenstoffatomen in der Alkylkette bedeutet, wobei die Ringkohlenstoffatome gegebenenfalls mit einem gerad- oder verzweigtkettigen Perfluoralkylrest mit 1 bis 12 Kohlenstoffatomen substituiert sein können, wobei $R_{\rm fl}$ und $R_{\rm f2}$ zusammengenommen insgesamt mindestens 3 Konlenstoffatome aufweisen;

R_B einen Perfluoralkylenrest mit 2 bis 6 Kohlenstoffatomen bedeutet, der gegebenenfalls mit einem geradoder verzweigtkettigen Perfluoralkylrest mit 1 bis 12 Kohlenstoffatomen oder einem Perfluorcycloalkylrest mit 4-7 Kohlenstoffatomen substituiert ist;

M ein Kation mit einer Wertigkeit n bedeutet und

n eine ganze Zahl von 1 bis 4 ist, wobei die an der positiven Elektrode gemessene Spannung größer als 3,0 V gegen Li/Li+ ist.

- Batterie nach Anspruch 1, wobei M Lithium ist.
- Batterie nach Anspruch 1, wobei das Salz Lithiumbis(perfluorethansulfonyl)imid ist.
 - 4. Batterie nach Anspruch 1, die außerdem ein Salz, ausgewählt aus Alkalimetall-, Erdalkalimetall-, Seltenerdenmetall- und Metallsalzen der Gruppe IIB und IIIB von Anionen, ausgewählt aus NO₃-, BF₄-, PF₆-, AsF₆-, CIO₄-, SbF₆-, R_fSO₃-, wobei R_f einen Perfluoralkylrest mit 2 bis einschließlich 12 Kohlenstoffatomen bedeutet, Bis(perfluormethylsulfonyl)imid;

Rn R2N-(CF2) - SO2-X-

$$Z = N - (CF_2)_{\overline{n_1}} SO_2 - X$$

wobei X- -O-, -N-SO₂R₁₃ oder

bedeutet; Z -CF₂-, -O-, >NR_{f8} oder -SF₄- bedeutet; R_{f1} und R_{f2} unabhängig voneinander -CF₃, -C_mF_{2m+1} oder -(CF₂)_q-SO₂-X M+ sind; R_{f3}, R_{f4} und R_{f5} unabhängig voneinander -CF₃, -C_mF_{2m+1}, -(CF₂)_q-SO₂-X M+,

$$N - (CF_2)_{\overline{n_1}}$$

oder

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$$Z = N - (CF_2)_{\overline{n_1}}$$

bedeutei

 R_{f8} -CF₃, -C_mF_{2m+1} oder -(CF₂)_q-SO₂-X-M+ darstellt; R_{f6} und R_{f7} unabhängig voneinander Perfluoralkylenreste der Formel -C_rF_{2r}- darstellen; n_1 1-4 ist; r 1-4 ist; r 1-12 ist und r 1-4 ist; Bisperfluoralkylsulfonylmethidanionen der Formel R_f -SO₂-C-(R)-SO₂-R_f', in der R_f und R_f ' unabhängig voneinander Perfluoralkylreste mit 1 bis einschließlich 12 Kohlenstoffatomen bedeuten und R H, Br, Cl, I, einen Alkylrest mit 1 bis einschließlich 20 Kohlenstoffatomen, einen Aryl- oder Alkylarylrest bedeutet und einem Tris(perfluoralkylsulfonyl)methidanion der Formel -C(SO₂R_f)(SO₂R_f'), wobei R_f , R_f ' und R_f * unabhängig voneinander Perfluoralkylreste mit 1 bis einschließlich 12 Kohlenstoffatomen bedeuten, umfaßt.

- 5. Batterie nach Anspruch 1, die außerdem LiNO3 umfaßt.
- Batterie nach Anspruch 1, wobei die Batterie wiederaufladbar ist.
- Batterie nach Anspruch 6, wobei die Batterie einen Lebenszyklus von mehr als 50 Zyklen besitzt.

Revendications

10 1. Pile comprenant

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au moins une électrode positive;

au moins une électrode négative ;

au moins un collecteur de courant en aluminium; et

une composition d'électrolyte comprenant un sel placé dans une matrice, ledit sel répondant à la formule

 $[(\mathsf{R}_{\mathsf{fl}}\,\mathsf{SO}_2)(\mathsf{R}_{\mathsf{f2}}\mathsf{SO}_2)\mathsf{N}]_{\mathsf{n}}\mathsf{M} \tag{I}$

OU

 $\begin{bmatrix} SO_2 \\ SO_2 \end{bmatrix} N$

dans laquelle

 \exists_n et \exists_2 représentent chacun indépendamment un groupe perfluoroalkyle linéaire ou ramifié contenant $\exists 1$ à 12 atomes de carbone, un groupe perfluorocycloalkyle ou un groupe perfluorocycloalkyleperfluoroalkyle contenant 4-7 atomes de carbone dans le cycle et de 1-4 atomes de carbone dans la chaîne alkylique, les atomes de carbone du cycle pouvant être éventuellement substitués par un groupe perfluoroalkyle linéaire ou ramifié contenant de 1 à 12 atomes de carbone, Rf_1 et Rf_2 , pris ensemble, contenant un total d'au moins 3 atomes de carbone;

Efg représente une partie perfluoroalkylène comprenant de 2 à 6 atomes de carbone, éventuellement substituée par un groupe perfluoroalkyle linéaire ou ramifié contenant 1 à 12 atomes de carbone ou par un groupe perfluorocycloalkyle contenant de 4 à 7 atomes de carbone;

M représente un cation avec une valence égale à n; et

n représente un nombre entier de 1 à 4;

cans laquelle la tension mesurée à l'électrode positive est supérieure à 3,0 volts vs Li/Li+.

- 2. Pile de la revendication 1, dans laquelle M représente le lithium.
- 3. Pile de la revendication 1, dans laquelle fedit sel est du bis(perfluoroéthanesulfonyl)imide de lithium.
 - 4. Pile de la revendication 1, comprenant en outre un sel choisi parmi des sels de métaux alcalins, de métaux alcalino-

terreux, d terres rares et de métaux des groupes IIB et IIIB d'anions choisis parmi NO₃°, BF₄°, PF₆°, AsF₆°, CIO₄°, SbF₆-, R₁SO₃- dans lequel R₁ représente un groupe perfluoroalkyle contenant de 2 à 12 atomes de carbone, inclus), bis(perfluorométhylsulfonyl)imide ; $R_{fl}R_{f2}N$ -(CF_2)_{n1}- SO_2 -X*;

dans laquelle X⁻ représente -O⁻, N⁻SO₂R_B, ou

 $(R_{f5}SO_2)-C-(SO_2R_{f4})$;

Z représente -CF₂-, -O-,

ou -SF₄- ; R_{f1} et R_{f2} représentent, indépendamment, -CF₃, -C_mF_{2m+1}, ou -(CF₂)_q-SO₂-X·M+ ; R_{f3} , R_{f4} et R_{f5} représentent sentent, indépendamment, -CF3, C_mF_{2m+1} , -(CF2)q-SO2-X-M+,

ou

 R_{f8} représente -CF₃, C_mF_{2m+1} , ou -(CF₂)_q-SO₂-X^{*}M⁺ ; R_{f6} et R_{f7} représentent, indépendamment, des parties perfluoroalkylène ayant pour formule -C_rF_{2r}-; n₁ vaut 1-4; r vaut 1-4; m vaut 1-12; et q vaut 1-4; des anions bisperfluoroalkylsulfonylméthide ayant pour formule $R_r SO_2 - C^-(R) - SO_2 - R_r$ dans laquelle R_f et R_f représentent, indépendent la company de la comp damment, des groupes perfluoroalkyle contenant entre 1 et 12 atomes de carbone, inclus, et R représente H, Br, CI, I, un groupe alkyle contenant entre 1 et 20 atomes de carbone, inclus, des groupes aryle ou alkylaryl; et un anion tris(perfluoroalkylsulfonyl)méthide de formule -C(SO₂R_f)(SO₂R_f) dans laquelle R_f, R_f et R_f représentent, indépendamment, des groupes perfluoroalkyle contenant de 1 à 12 atomes de carbone, inclus.

- 5. Pile de la revendication 1, comprenant en outre du LiNO₃.
- Pile de la revendication 1, dans laquelle la pile est recharg able.

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7. Pile de la revendication 6, la pile ayant une durée de vie en cycles supérieure à 50 cycles.

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